TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN 1—Continued

21 CFR Section; activity	Number of respondents	Number of responses per respondent	Total annual responses	Average burden per response	Total hours
607.22(b); Written waiver request	1	1	1	1	1
Total					1,460

¹There are no capital costs or operating and maintenance costs associated with this collection of information.

Based on our evaluation of calendar year 2022 data from CBER's Blood Establishment Registration and Product Listing system, we have adjusted the currently approved burden estimate we attribute to establishment registration and product listing to reflect a decrease in product listing updates and an increase in the number of initial registrations. Our estimated burden for the information collection reflects an overall decrease of 36 hours.

Dated: March 6, 2024.

Lauren K. Roth,

Associate Commissioner for Policy.
[FR Doc. 2024–05215 Filed 3–11–24; 8:45 am]
BILLING CODE 4164–01–P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Food and Drug Administration

[Docket No. FDA-2023-N-3847]

Agency Information Collection Activities; Submission for Office of Management and Budget Review; Comment Request; Adverse Experience Reporting for Licensed Biological Products; and General Records

AGENCY: Food and Drug Administration, HHS.

ACTION: Notice.

SUMMARY: The Food and Drug Administration (FDA, Agency, or we) is announcing that a proposed collection of information has been submitted to the Office of Management and Budget (OMB) for review and clearance under the Paperwork Reduction Act of 1995. DATES: Submit written comments (including recommendations) on the collection of information by April 11,

ADDRESSES: To ensure that comments on the information collection are received, OMB recommends that written comments be submitted to https://www.reginfo.gov/public/do/PRAMain. Find this particular information collection by selecting "Currently under Review—Open for Public Comments" or by using the search function. The OMB

control number for this information collection is 0910–0308. Also include the FDA docket number found in brackets in the heading of this document.

FOR FURTHER INFORMATION CONTACT:

Domini Bean, Office of Operations, Food and Drug Administration, Three White Flint North, 10A–12M, 11601 Landsdown St., North Bethesda, MD 20852, 301–796–5733, *PRAStaff@fda.hhs.gov.*

SUPPLEMENTARY INFORMATION: In compliance with 44 U.S.C. 3507, FDA has submitted the following proposed collection of information to OMB for review and clearance.

Adverse Experience Reporting for Licensed Biological Products; and General Records—21 CFR Part 600

OMB Control Number 0910–0308— Extension

This information collection helps support implementation of statutory and regulatory authorities that govern adverse experience reporting. Under the Public Health Service Act (PHS Act) (42 U.S.C. 262), FDA may only approve a biologics license application for a biological product that is safe, pure, and potent. When a biological product is approved and enters the market, the product is introduced to a larger patient population in settings different from clinical trials. New information generated during the postmarketing period offers further insight into the benefits and risks of the product, and evaluation of this information is important to ensure its safe use. Regulations implementing adverse experience reporting (AER) requirements applicable to biological products are codified in part 600 (21 CFR part 600). Regulations applicable to combination products subject to regulations in part 600 are found in part 4 (21 CFR part 4)—Regulation of Combination Products. The collections of information are intended to enable FDA to take actions necessary for the protection of the public health in response to reports of adverse experiences related to biologics licensed

under any provision of section 351 of the PHS Act.

To assist respondents with the reporting provisions of the information collection, FDA has created both paperbased and electronic forms. Information may be submitted electronically through MEDWATCH or the Vaccine Adverse Experience Reporting System (VAERS). AER reports are filed using the MEDWATCH Form FDA-3500A (approved under OMB control numbers 0910-0291 and 0910-0645) or the VAERS-1. Both versions of the forms and instructions are available from the internet at https://vaers.hhs.gov. The forms may also be downloaded, completed, and submitted to the Agency by mail or facsimile.

For operational efficiency, on March 20, 2023, we requested, and OMB has approved, the addition of burden attributable to provisions set forth in part 4, subpart B, previously included in OMB control number 0910–0834. When information regarding an event that involves a death or serious injury, or an adverse event, associated with the use of the combination product is received by the product sponsor, the information must be provided to the other constituent part applicant(s) no later than 5 calendar days after receipt under § 4.103. Relatedly, § 4.104 explains how and where to submit reports.

In the **Federal Register** of September 28, 2023 (88 FR 66856), we published a 60-day notice requesting public comment on the proposed collection of information. We received one comment regarding our estimate of 28 hours per response for periodic adverse experience reports. The comment suggested we lower that estimate but provided no data or explanation in support of the proposed reduction. While we have therefore made no adjustment in our burden estimate, we encourage further comment regarding a basis for assessing burden for the scope of information collection activity covered by the applicable regulations and associated forms.

Respondents: Respondents to this collection of information are manufacturers of biological products (including blood and blood

components) and any person whose name appears on the label of a licensed biological product. We estimate the burden of this collection of information as follows:

TABLE 1—ESTIMATED ANNUAL REPORTING BURDEN—BIOLOGICAL PRODUCTS 1

21 CFR Section; activity	Number of respondents	Number of responses per respondent	Total annual responses	Average burden per response (in hours)	Total hours
600.80(c)(1), 600.80(d), and 600.80(e); postmarketing 15-day Alert Reports 600.82; notification of discontinuance or interruption in manufacturing	109 23 109 172	3,806.95 1.435 3,697 5.727	414,958 33 402,973 985	1 2 28 1	414,958 66 11,283,244 985
600.80(h)(2), 600.81(b)(2), and 600.90; waiver requests	35	1.886	66		11,699,319

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

TABLE 2—ESTIMATED ANNUAL REPORTING BURDEN—BIOLOGICAL PRODUCTS 1

21 CFR Section; activity	Number of recordkeepers	Numbers of records per recordkeeper	Total annual records	Average burden per recordkeeper (in hours)	Total hours
600.122; Maintenance of Records	131 216	40.145 3.4028	5,259 735	32 24	168,288 17.640
600.12(b)(2); Recall Records	109	7,503.95	817,931	1	817,931
Total					1,003,859

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

² The recordkeeping requirements in § 610.18(b) are included in the estimate for § 600.12.

TABLE 3—ESTIMATED ANNUAL REPORTING BURDEN—COMBINATION PRODUCTS 1

21 CFR Section; activity	Number of respondents	Number of disclosures per respondents	Total annual disclosures	Average burden per disclosure (in hours)	Total hours
4.102, 4.103, 4.104, 4.105; Postmarketing Safety Reporting for Combination Products, including associated reports and sharing information with other constituent part applicants.	11	18	198	0.35 (21 minutes)	69

¹ There are no capital costs or operating and maintenance costs associated with this collection of information.

The burden for this information collection has changed since the last OMB approval. The reporting and recordkeeping burden has increased mostly due to an increase in the number of AER reports submitted to FDA and the associated recordkeeping with these reports. We have also added burden we believe attributable to post marketing safety reporting and attendant recordkeeping and disclosures, as required under part 4, subpart B.

Dated: March 7, 2024.

Lauren K. Roth,

 $Associate\ Commissioner\ for\ Policy.$ [FR Doc. 2024–05222 Filed 3–11–24; 8:45 am]

BILLING CODE 4164-01-P

DEPARTMENT OF HEALTH AND HUMAN SERVICES

Meeting of the Presidential Advisory Council on HIV/AIDS

AGENCY: Department of Health and Human Services, Office of the Secretary,

Office of the Assistant Secretary for Health.

ACTION: Notice of a meeting.

SUMMARY: As stipulated by the Federal Advisory Committee Act, the U.S. Department of Health and Human Service is hereby giving notice that the Presidential Advisory Council on HIV/ AIDS (PACHA or the Council) will convene the 80th full council meeting on Wednesday, March 27-Thursday, March 28, 2024. The meeting will be open to the public and there will be a public comment session during the meeting; pre-registration is required to provide public comment. To pre-register to provide public comment, please send an email to PACHA@hhs.gov and include your name, organization, and title by close of business Monday, March 18, 2024. If you decide you would like to provide public comment but do not pre-register, you may submit your written statement by emailing PACHA@hhs.gov by close of business

Thursday, April 4, 2024. The meeting agenda will be posted on the PACHA page on *HIV.gov* at *https://www.hiv.gov/federal-response/pacha/about-pacha* prior to the meeting.

DATES: The meeting will convene on Wednesday, March 27, 2024 from approximately 10:00 a.m. (Eastern) –7:00 p.m. (Eastern) and Thursday, March 28, 2024 from approximately 10:00 a.m. (Eastern) to 4:45 p.m. (Eastern).

ADDRESSES: Texas Southern University, 3100 Cleburne Avenue, Houston, TX 77004. To attend the meeting virtually, please visit *www.hhs.gov/live*.

FOR FURTHER INFORMATION CONTACT: Ms. Caroline Talev, MPA, Senior Management Analyst, at *PACHA@hhs.gov* or *Caroline.Talev@hhs.gov*. Additional information can be obtained by accessing the Council's page on the *HIV.gov* site at *www.hiv.gov/pacha*.

SUPPLEMENTARY INFORMATION: PACHA was established by Executive Order 12963, dated June 14, 1995, as amended